

The male and female perspective in the link between male infant care and mating behaviour in Barbary macaques

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Title: The male and female perspective in the link between male infant care and mating behaviour in Barbary macaques

Short title: Infant care and mating in macaques

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ABSTRACT

Infant care from adult males is unexpected in species with high paternity uncertainty. Still, males of several polygynandrous primates engage in frequent affiliative interactions with infants. Two non-exclusive hypotheses link male infant care to male mating strategies. The paternal investment hypothesis views infant care as a male strategy to maximise the survival of sired offspring, while the mating effort hypothesis predicts that females reward males who cared for their infant by preferably mating with them. The paternal investment hypothesis predicts a positive relationship between the distribution of matings and subsequent infant care, whereas the mating effort hypothesis between care and subsequent matings. Both hypotheses are usually tested from the female perspective, – based on the proportion of a female's or her infant's interactions with a specific male relative to her or her infant's interactions with all males; but not from the male perspective, based on the proportion of a male's interactions with a specific female relative to his interactions with all females.

We tested the relationships between care and mating from both perspectives in Barbary macaques. Mating predicted subsequent care and care predicted subsequent mating when viewed from the male but not the female perspective. Males mainly cared for infants of their main mating partners, but infants were not mainly cared for by their likely father. Males mated more with the mothers of their favourite infants, but females did not mate more with the main caretakers of their infants.

We suggest that females do not choose their mating partners based on previous infant care, increasing paternity confusion. Males might try to increase paternal investment by distributing the care according to their own instead of female mating history. Further, males pursue females for mating opportunities based on previous care.

Keywords: Male infant care – infant handling – paternal investment – mating effort –

Macaca sylvanus

INTRODUCTION

Mammalian males and females differ in their reproductive strategies. Females are limited in the number of offspring they can produce by the time and energetic costs of gestation and lactation (Geary, 2015). Males are limited by the number of mating opportunities they can secure and face paternity uncertainty (Trivers, 1972). These differences lead to conflict between females and males, in which the two sexes use different strategies to increase their own fitness, sometimes constraining the fitness of the opposite sex (Gavrilets, Arnqvist, & Friberg, 2001; Parker, 1979). The typical example of such a sexual conflict is the interplay between male infanticide, proposed to be an adaptive male strategy (Hrdy, 1979; van Schaik, Pradhan, & van Noordwijk, 2004), and paternity confusion, a female counter-strategy against infanticide. However, sexual conflict is expressed by various behavioural strategies, including the number of mating partners, mating frequency and its timing, maintenance of intersexual relationships and relative parental investment (Stumpf, Martinez-Mota, Milich, Righini, & Shattuck, 2011).

Postnatal maternal investment in the form of lactation is ubiquitous in mammals and typically extends to other forms of care-giving (Clutton-Brock & Parker, 1992; Trivers, 1972). The degree of male paternal investment varies strongly across taxa. In the majority of mammalian species mothers are the sole caretakers, but in some species (e.g. some carnivores, rodents and primates; Woodroffe & Vincent, 1994) males actively take care of infants. Sexual selection theory links the variation in male investment with paternity certainty (Trivers 1972). Males may forfeit mating opportunities when

89 taking care of infants and, when paternity is uncertain, risk misdirecting paternal
90 investment to the offspring of other males (Clutton-Brock & Parker, 1992; Andreas Paul,
91 Preuschoft, & van Schaik, 2000; Trivers, 1972). It is thus expected that the higher the
92 paternity uncertainty, the less likely males are to invest into infants.

93 Although the risk of misdirected paternal investment is likely to be high in
94 polygynandrous primates, males in some species do associate closely with infants
95 (Huchard et al., 2012), groom and carry them (Deag, 1980; Estrada & Sandoval, 1977),
96 support them in conflicts (Buchan, Alberts, Silk, Altmann, & others, 2003; Minge,
97 Berghänel, Schülke, & Ostner, 2016) and protect them from harassment and infanticide
98 (Palombit, Seyfarth, & Cheney, 1997; Paul, Preuschoft, & van Schaik, 2000). Two
99 hypotheses explain such behaviour as a part of an adaptive male reproductive strategy.
100 The paternal investment hypothesis (Trivers, 1972) proposes that males are able to assess
101 their chances of paternity and direct their infant care accordingly, even in polygynandrous
102 species. It suggests that the probability of a male to care for an infant increases with his
103 chances of having sired the infant. Supporting this hypothesis, in yellow (*Papio*
104 *cynocephalus*; Altmann, 2001) and olive baboons (*P. anubis*; Smuts, 1985) males were
105 more likely to take care of infants if they had consorted (see Manson, 1997 for the
106 definition) or mated with an infant's mother in the preceding mating season than if they
107 did not consort/mate with her (see also Smuts & Gubernick, 1992). In chacma baboons
108 (*P. ursinus*), male-infant associations were predicted by consortship activities and/or
109 social bonds between males and females (Buchan et al., 2003; Moscovice et al., 2010),
110 that predicted paternity (Buchan et al. 2003). In crested macaques (*Macaca nigra*), the
111 likelihood of male-infant affiliation was significantly higher if the male was present in
112 the group at the infant's conception (Kerhoas et al., 2016). In Assamese (*M. assamensis*;

Ostner, Vigilant, Bhagavatula, Franz, & Schülke, 2013) and rhesus macaques (*M. mulatta*; Langos, Kulik, Mundry, & Widdig, 2013) the distribution of copulations predicted the distribution of subsequent male infant care, suggesting that males estimate their paternity chances when distributing infant care.

The mating effort hypothesis (Seyfarth, 1978) views male infant care as a reproductive strategy to trade investment into infants for future mating opportunities with the infant's mother, who monitors male care and distributes subsequent mating accordingly (Seyfarth 1978; Smuts 1985). It predicts a positive relationship between care and subsequent mating (Ménard et al., 2001; Seyfarth, 1978; Smuts, 1985). Evidence of this pattern is rather weak in baboons (Nguyen, Van Horn, Alberts, & Altmann, 2009; Weingrill, 2000) however, some findings supported this hypothesis in macaques. In Tibetan macaques (*M. thibetana*), the amount of infant care that a male provided predicted the number of females who later consorted with that male (Zhao, 1996). In Barbary macaques (*M. sylvanus*), males were likely to become the primary sexual partners of those females whose offspring they had previously cared for the most (Ménard et al. 2001), supporting the mating effort hypothesis. However, an earlier study on the same species did not find this pattern (Paul, Kuester, & Arnemann, 1996). Indirect evidence might also come from mountain gorillas (*Gorilla beringei beringei*), where those males that were most involved in infant care had the highest lifetime fitness, although they did not direct care specifically to their own offspring (Rosenbaum, Vigilant, Kuzawa, & Stoinski, 2018).

Both the paternal investment and mating effort hypotheses predict a link between the distributions of infant care and mating. The paternal investment hypothesis predicts a

positive relationship between the distribution of current mating and infant care during the subsequent birth season, whereas the mating effort hypothesis predicts a positive relationship between current infant care and subsequent mating. Notably, both patterns have been tested so far from the female perspective, that is based on the proportion of a female's or her infant's interactions with a specific male, relative to her/her infant's interactions with all males. Little attention has been paid to the perspective of males, i.e. the proportion of a male's interactions with a specific female/her infant, relative to his interactions with all females/her infants.

Although this difference may seem negligible, the two perspectives in fact provide different information. First, when testing whether mating predicts subsequent care, the female perspective is suitable to test the prediction of the paternal investment hypothesis, i.e. that males take into account the distribution of female matings across all males when distributing their infant care, because the probability of paternity is likely based on how often females mated with different males. However, the female perspective implies that males are able to monitor all matings in a group (i.e. use knowledge of third party interactions) which may not be the case in a promiscuous species. If males can only access their own mating history, a more parsimonious prediction is that males distribute care based on their own proportional mating with a specific female. To test this prediction, the link between mating and subsequent care needs to be investigated from the male perspective. Second, when testing whether care predicts subsequent mating, the female perspective is in accord with the original definition of the mating effort hypothesis, which assumes that females control the distribution of mating and prioritizes mating with those males that care the most for her infant (Seyfarth, 1978). However, the male perspective is needed to address whether males also preferentially pursue females (for mating

opportunities) on the basis of infant care, possibly indicating that the distribution of male infant care is influenced by a male's effort to establish a long-term relationship with a female. In summary, testing the relationship between mating and male care from both the female and male perspectives may help to understand male care in the context of male and female reproductive effort and tap into the question of the importance of individual (own) and social (the other's perspective) information for reproductive strategies.

Barbary macaque reproduction is characterized by high seasonality (with mating mostly concentrated into a three month period), a high degree of female ovarian cycle synchrony (Brauch et al., 2008; Küster & Paul, 1984; Taub, 1980b; Young et al., 2013) and promiscuity (Small 1990), all of which is likely to increase paternity confusion. Copulations are brief, are rarely non-ejaculatory and males rarely achieve more than one copulation during consociation (Taub 1982). A consociation is often terminated by females who mate with multiple males at rapid succession (Small, 1990). Males rarely exhibit mate-guarding and male reproductive skew is low (Bissonnette et al., 2011), which is possibly due to the inability of males to monopolize females. Females show cyclic changes in sexual behavior and sexual swellings (Brauch et al., 2007; Young et al., 2013; but see Small, 1990), but they also display post-oestrous swelling which is thought to increase paternity uncertainty (Young et al., 2013). Indeed, males also mate with females when the probability of the conception is low, including during post-conception periods (Young et al., 2013). Finally, females produce copulation calls which may function to induce male-male competition and lead to more effective paternity confusion (Pfefferle, Brauch, Heistermann, Hodges, & Fischer, 2008; Semple, 1998).

Although these characteristics likely increase paternity confusion, Barbary macaque males frequently interact with infants (Whitten, 1987). Interactions between males and infants can either take the form of dyadic interactions between a single male and an infant during which a male carries, cradles, and/or grooms an infant, or take the form of triadic male-infant-male interactions, during which two males jointly manipulate an infant (Taub, 1980b). Both interaction types are unevenly distributed among males as well as infants (Deag, 1980; Kubenova et al., 2017; Ménard et al., 2001; Taub, 1984). Triadic male-infant-male interactions in macaques are proposed to primarily serve a social function among males (Deag & Crook, 1971; Kalbitz, Schülke, & Ostner, 2017; Paul et al., 1996), but explanations for dyadic male-infant interactions remain inconclusive (e.g. Paul et al., 1996; Taub, 1980b). Previous studies on Barbary macaques have shown that males are the primary caretakers of neither their genetic offspring (Ménard et al., 2001; Ménard, Scheffrahn, Vallet, Zidane, & Reber, 1992; Paul, Kuester, & Arenmann, 1992) nor their likely offspring based on previous mating success (Paul et al. 1996), speaking against the paternal investment hypothesis. Instead, one study has shown that males are likely to become the primary sexual partners of those females whose offspring they had previously cared for the most (Ménard et al., 2001), supporting the mating effort hypothesis. The conclusions are based on the link between the care and mating viewed from the female perspective. The male perspective has not been investigated.

We investigated both the female and the male perspectives of the relationships between mating and subsequent care and between care and subsequent mating in Barbary macaques. This resulted into four predictions (Table 1):

- 1) We tested the paternal investment hypothesis from the female perspective to assess whether males monitor female matings and preferentially care for infants who are likely to be their offspring, based on the male's mating history with a specific female. To this end we examined the relationship between mating and subsequent care, predicting that the more a female mated with a specific male (relative to all her matings) the more care her infant will subsequently receive from this male (relative to all male care the infant receives).
- 2) We tested the paternal investment hypothesis from the male perspective to assess whether males monitor their own previous matings when distributing infant care. We examined the relationship between mating and subsequent care, predicting that the more a male mated with a specific female (relative to all his matings), the more he will subsequently care of her infant (relative to all his infant care).
- 3) We tested the mating effort hypothesis from the female perspective to assess whether females reward males who cared for their infant by preferably mating with them. We examined the relationship between care and subsequent mating, predicting that the more care an infant received from a specific male (relative to all male care the infant received), the more the infant's mother will mate with him (relative to all her matings).
- 4) We tested the mating effort hypothesis from the male perspective to assess whether males preferentially pursue mothers of their preferred infants for mating opportunities. We examined the relationship between care and subsequent mating, predicting that the more a male cared for a specific infant (relative to all infant care from him), the more he will subsequently mate with the infant's mother (relative to all his matings).

229 *Table 1 placed here*

230 **METHODS**

231 Data collection

232 We conducted this study in Ifrane National Park in the Middle Atlas Mountains
233 of Morocco (33-240°N, 005-120°W) under the research permission (No. 253/2013,
234 26/2014) of the Haut Commissariat aux Eaux et Forêts et à la Lutte Contre la
235 Désertification of Morocco. The research complied with international, national and/or
236 institutional guidelines for the ethical treatment of primates and with Directive
237 2010/63/EU. We adhered to the legal requirements of Morocco and ASAB/ABS
238 Guidelines for the care and use of animals. We followed one group (Green Group) of
239 macaques, which was well habituated to the presence of human observers. Focal
240 continuous observations (Altmann, 1974) of infants were conducted by BK over two field
241 seasons, including two consecutive birth seasons (April – August 2013, April – June
242 2014). All individuals including infants were individually recognized based on their
243 physical features (facial features, specific coloration patterns on head when changing the
244 fur for infants).

245 In both seasons the group consisted of 6 adult males (> 5 years), 1 subadult (5
246 years) male, and 6 adult females (> 5 years). There were nineteen juveniles in 2013, and
247 twenty in 2014. All adult females gave birth in both seasons, resulting in six infants each
248 season (5 females, 1 male, born between April 5 and 30 in 2013 and 3 females, 3 males,
249 born between April 10 and 27 in 2014). Infants were followed for 2-hour observation
250 sessions, during which all social interactions between the focal infant and adult males
251 were recorded. We pseudorandomized the order in which infants were observed to ensure

that they were all observed equally often at different times of day. In both seasons, we started data collection once four infants were born. Data collection on the twelve infants yielded a total of 903 hours of observation, 582 hours in 2013 (between 83 and 109 hours per infant, median = 101), and 321 hours in 2014 (between 44 and 61 hours per infant, median = 55).

We recorded all interactions between the focal infant and adult male(s), indicative of male care or affiliation including the start and termination of proximity (within 1.5 distance) as well as start and termination of active male care and “passive” body contact (see below). The start of proximity was recorded when a male approached the infant (to within a 1.5 m radial distance), when an infant approached a male, or was brought to a male by another individual. The end of proximity was recorded when the infant left the male (or vice versa), or when the infant was carried away by another individual. Active male care included carrying, cradling and grooming; passive body contact was recorded once an infant was in body contact with a male without apparent involvement of a male (e.g. infant resting, crawling in body contact with a male or crawling over his body). Triadic male-infant-male interactions were not included in any of those categories, as they may be motivated by factors unrelated to our main question, e.g. male relationship management (Deag, 1980; Paul et al., 1996).

Mating behaviour and dyadic dominance-submission interactions were recorded *ad libitum* (Altmann, 1974) on adult macaques by several researchers between April 2013 and September 2014. All matings were recorded between October 2013 and February 2014 (further referred to as the mating season).

Data analysis

To assess the distribution of male care for infants, we used a composite sociality index (CSI; Silk, Altmann, & Alberts, 2006). We used five categories of affiliative dyadic interactions (recorded during focal observations and controlled for dyadic focal observation time) between males and infants to calculate the CSI, which were all positively correlated (row-wise matrix correlation, average row-wise tau: 0.68-0.76). These behavioural categories were (1) frequency and (2) duration of active male care, (3) duration of passive body contact, (4) frequency of male approaches into infant's 1.5m proximity and (5) duration of time spent within 1.5 m proximity (excluding duration of passive body contact and active male care).

We calculated two different CSIs, one for infants (CSII) and one for males (CSIm), in order to express the distribution of care from the infant's (corresponding with the female's) perspective and male's perspective, respectively.

To generate CSII, we divided the value for each of the five behavioural categories (Beh. infant-male) between the male and the infant by the average value across all dyads this infant formed with all males ($\overline{\text{Beh. infant - all males}}$) divided by the number of behavioural categories, i.e. five, following the formula

$$CSII = \frac{\sum_{i=1}^5 \text{Beh. infant - male} / \overline{\text{Beh. infant - all males}}}{5}.$$

This quantified how much care the infant received from a particular male relative to all other males.

To generate CSIm, we divided the value for each of the five behavioural categories between the male and the infant (Beh. infant-male) by the average across all dyads this

male formed with all infants ($\overline{\text{Beh. male} - \text{all infants}}$) divided by the number of categories, following the formula

$$CSIm = \frac{\sum_{i=1}^5 \text{Beh. infant} - \text{male} / \overline{\text{Beh. male} - \text{all infants}}}{5}.$$

This quantified how much a particular male cared for a particular infant relative to all other infants. We assessed CSIs for both perspectives separately for birth season 2013 and 2014.

To assess mating distribution from the female perspective (female matings), we calculated the proportion of matings each male achieved with a female from all matings of this female (with all males). To assess mating distribution from the male perspective (male matings), we calculated the proportion of a male's matings with each female from all his matings (with all females). Values could range between 0 and 1 (0 = no matings with given partner, 1 = all matings with given partner only). We run row-wise matrix correlation (using Matman 1.1.4 program; Netto, Hanegraaf, & De Vries, 1993) to compare female and male matings.

To assess dominance rank, we entered dominance-submission interactions (recorded as *Ad libitum*) into a winner-loser matrix and built a hierarchy based on the standardized normalized David's score (De Vries, Stevens, & Vervaecke, 2006). The hierarchy was assessed separately for males and females, and birth season 2013 and birth season 2014. Dominance rank was based on 213 interactions in females and 45 in males in 2013, and 249 in females and 71 in males in 2014, respectively.

Statistical analysis

We used linear mixed models (LMMs) using the lme4 package (Bates, Maechler, Bolker, Walker, & others, 2014) in R 3.1.1 (R Core Team, 2014). To evaluate the effects of the predictors we computed the 95% confidence intervals (using the *confint* function in lme4). We constructed a separate model for each of four predictions. In all four models, female/infant and male IDs were entered as random effects. Every possible male-female, respectively male-infant combination (N=36) entered the analysis as an independent data point. Depending on the perspective (female versus male), we controlled for the male, respectively female dominance rank.

RESULTS

All infants were observed in 1.5 m proximity with all males and in active care or passive body contact with 0-6 males (median=6) over the respective birth seasons. Infants spent between 0% and 15% of observation time in active care (mean=3.6 ± SD 5.3%) and between 0% and 4% in passive body contact (mean=5.2 ± SD 6.1%). Only one infant was never observed in active care or passive body contact with any male. From 36 possible male-infant dyads, 18 (50%) dyads were observed in active care and/or passive body contact in 2013 and 25 (69%) dyads in 2014. CSIs were calculated based on 5,829 interactions in period 2013 (range per infant = 221-2,206; mean=972 ± SD 937) and 4,529 interactions in 2014 (range per infant = 404-1,525, mean=755 ± SD 421). CSI values ranged between 0.027 and 4.150 for the infant perspective (median=0.471) and between 0.007 and 4.602 for the male perspective (median=0.379).

We recorded a total of 152 matings (range per female=6-49; mean=25.33 ± SD 16.27; range per male=14-50; mean=25.33 ± SD 15.04). All females and males had at least five mating partners from a possible six. The proportion of female matings with

different males ranged between 0 and 0.50 (median=0.14). The proportion of male matings with different females ranged between 0 and 0.58 (median=0.10). Proportions of female and male matings significantly correlated (Kr test, row-wise tau=0.461, $p<001$). Further details about the distributions of CSIs and matings are provided as supporting information in the electronic supplement.

Model 1: Paternal investment hypothesis, female perspective

The care an infant received from a male, relative to all male care that infant received from any male was neither predicted by how often, in the previous mating season, the infant's mother had mated with this male relative to all her matings, (LMM: Estimate = -0.623; SE = 1.778; $t = -0.351$; CI95% = -4.050, 2.803) nor by male dominance rank (GLMM: Estimate = 0.021; SE = 0.043; $t = 0.484$; CI95% = -0.063, 2.803).

Model 2: Paternal investment hypothesis, male perspective

The care a male directed to a specific infant, relative to all the care provided by that male, was predicted by how often, in the previous mating season, that male had mated with the infant's mother relative to all his matings (LMM: Estimate = 3.949; SE = 1.401; CI95% = 1.248, 6.651) and was not affected by female dominance rank (GLMM: Estimate = 0.019; SE = 0.026; CI95% = -0.031, 0.069).

Figure 1 placed here

Model 3: Mating effort hypothesis, female perspective

How often a female mated with a specific male relative to all her matings, was neither predicted by the amount of care her infant had received from the male relative to all care the infant had received, in the previous birth season (LMM: Estimate = 0.014; SE

= 0.016; CI95% = -0.092; 0.045), nor by male rank (LMM: Estimate = 0.011; SE = 0.006; CI95% = -4.809; 0.021).

Model 4: Mating effort hypothesis, male perspective

The proportion of male's matings with a female, relative to all his matings, was predicted by how much care that male had directed to the female's infant (relative to all his infant care), in the previous birth season (LMM: Estimate = 0.039; SE = 0.018; CI95% = 0.003; 0.074), but was not predicted by female rank (LMM: Estimate = 0.003; SE = 0.0004; CI95% = -0.004; 0.010).

Figure 2 placed here

DISCUSSION

We investigated in wild Barbary macaques the relationship between mating and subsequent care, and between care and subsequent mating both from the female and the male perspective to test two non-exclusive hypotheses, the paternal investment and the mating effort hypothesis. When tested from the female perspective, neither hypotheses were supported; conversely both the paternal investment and the mating effort hypotheses were supported when tested from the male perspective.

Model 1: Paternal investment hypothesis, female perspective

The paternal investment hypothesis predicts that males care preferentially for their own offspring. Based on genetic data there is evidence for this pattern in some (e.g., yellow baboons: Buchan et al., 2003; Onyango, Gesquiere, Altmann, & Alberts, 2013; chacma baboons: Huchard et al., 2012; rhesus macaques: Langos et al., 2013; Assamese

macaques: Ostner et al., 2013), but not in other species (Kerhoas et al., 2016; Paul et al., 1996). A number of behavioural studies (rhesus macaques: Berenstein, Rodman, & Smith, 1981; Assamese macaques: Ostner et al., 2013; chacma baboons: Moscovice et al., 2010) suggest that males may distribute care depending on their previous mating history, potentially estimating their paternity chances (per infant) and adjusting care levels accordingly. This adjustment based on past mating history was neither confirmed in previous studies on Barbary macaques, where the mother's main sexual partner did not become the infant's main caretaker (Ménard et al., 2001; Paul et al., 1996), nor in the present study.

While this result does not align with the paternal investment hypothesis, we cannot conclusively reject it. Mating frequency may not be positively related to an increased probability of siring an infant (Ménard et al. 2001). Given that genetic data on paternity were not available in our study, we cannot rule out that genetic fathers did indeed become the main caretakers of their offspring despite having mated less than other males with the infant's mother. In this case, males may base their approximation of siring success on information we did not record, for example the timing of mating in relation to conception probability (Brauch et al., 2007; Semple & McComb, 2000; Young, Majolo, Heistermann, Schülke, & Ostner, 2013). It is unclear whether Barbary macaque males are able to estimate the timing of conception based on female cues. Although the size of sexual swellings may serve as a reliable signal of female fertility (Brauch et al., 2007; Young et al., 2013; but see Small, 1990), males also mate with females when the probability of the conception is low, and even during post-conception periods (Young et al., 2013). Further, females in our study mated in rapid succession with up to four different

males within one day. Thus, even if males could have estimated and considered the reproductive state of their mating partners, paternity uncertainty likely remained high.

Model 2: Paternal investment hypothesis, male perspective

Female promiscuity may make it difficult for males to keep track of female matings with other males. This might be why our and other studies (Ménard et al. 2001; Paul et al. 1996) do not support the paternal investment hypothesis when tested from the female perspective. Instead, we found that males cared more for the infants of their most frequent mating partners, supporting the paternal investment hypothesis from the male perspective. It is possible, that under the conditions of female promiscuity and incomplete fertility information, a male's best strategy to increase paternal investment might be to track their own matings and base care pattern on their own mating history. The crucial difference here is between private versus third party knowledge; only if the information males have on female mating activity with any male is sufficiently reliable, males can match their subsequent infant care accordingly. If such a third party information is noisy, males may use their own matings across females as a proxy. In our study, the proportions of female and male matings were correlated, thus tracking of one's own mating history (in the absence of more reliable information about paternity), may result in an adaptive benefit in terms of a decreased probability of directing care to a non-related infant. However, this benefit may remain obscure in the results due to our small sample size and lack of paternity data, which is a limitation of our study. However, if males make mistakes when estimating their paternity, we cannot exclude that even with genetic data, the relationship between care and paternity might be ambiguous in studies with small sample sizes.

An alternative, non-mutually exclusive driver of male care, particularly in Barbary macaques, is improved access to infants for frequent triadic male-infant-male interactions. These interactions help males to establish and maintain social bonds with other males (Kalbitz et al. 2017; Kuběnová et al., 2019; Kümmerli & Martin, 2008; Paul et al., 1996) which may reduce aggression among males (Deag & Crook, 1971) and help them to retain support in coalition (Paul et al., 1996). The benefit of social bonding may outweigh the cost of infant care. Concentrated in birth season, infant care in Barbary macaques does not represent a significant cost in terms of lost mating opportunities. The high body weight ratio between males and infants (Fooden, 2007) also suggests relatively low cost in terms of energy investment (Woodroffe & Vincent, 1994; Wright, 1990).

3. Mating effort hypothesis, female perspective

In contrast to the paternal investment hypothesis, the mating effort hypothesis proposes that males benefit from infant care if it increases their subsequent mating success with the infants' mothers (Seyfarth, 1978; Smuts, 1985; Smuts & Gubernick, 1992). That is, if a male provides an infant with care the mother may subsequently provide that male with mating privileges (Hector, Seyfarth, & Raleigh, 1989; Small, 1989). Consequently, the relationship between care and subsequent mating should be most apparent in species with a high potential for female mate choice (Seyfarth, 1978; Smuts, 1985). Female Barbary macaques are very active in sexual interactions (Small, 1990), initiating and terminating consortship (Taub, 1980a, p.292), and using copulations calls to manipulate paternity (Pfefferle et al., 2008; Semple, 1998). In line with this active sexual role of females, the mating effort hypothesis from the female perspective was supported in a previous study on Barbary macaques, showing that females preferentially mated with males that had cared for their previous infants (Ménard et al., 2001). In contrast, our data

as well as results of yet another study on Barbary macaques (Paul et al., 1996), did not provide evidence for the mating effort hypothesis as females did not adjust their mating rate depending on the amount of care their infants had received from specific males prior to the mating season. It is possible that the female active role in sexual behaviour does not reflect mate choice but aids females to mate with as many males as possible to increase paternity confusion (Small, 1990; Taub, 1980a). The importance of paternity confusion has been linked to the degree of the risk of infanticide by males (Hrdy, 1979; van Noordwijk & van Schaik 2000). In Barbary macaques, females might benefit from paternity confusion by reducing the risk of infanticide (Hrdy, 1979; van Noordwijk & van Schaik 2000) and also by securing male care for their offspring from several males (Taub, 1980a but see also Small, 1990).

Model 4: Mating effort hypothesis, male perspective

From the male perspective, the relationship between infant care and subsequent mating holds; males were more likely to mate with those females whose infants they cared for more in the previous birth season. This does not support the original formulation of the mating effort hypothesis, which proposes increased male mating success based on previous infant care from the female perspective. Instead, it is consistent with the “friends with benefits hypothesis” (Ostner et al., 2013), which relates mating patterns to stable affiliative relationships between the sexes. The influence of male-female affiliative relationships on the distribution of male care for infants has been demonstrated in chimpanzees (Langergraber, Mitani, Watts, & Vigilant, 2013), and several species of baboons (Goffe, Zinner, & Fischer, 2016; Moscovice et al., 2010; Palombit et al., 1997; Städele et al., 2019; Weingrill, 2000) and macaques (Aureli & Yates, 2010; Haunhorst,

Schülke, & Ostner, 2016; Hill, 1990; Kerhoas et al., 2016; Kulik, Amici, Langos, & Widdig, 2015; Massen et al., 2012; Ostner et al., 2013), but not yet in Barbary macaques (Small, 1990). It is possible that males establish affiliative relationships with females with whom they also preferentially mate. Females tolerate those males, allowing them to mate with them and to interact with their infants. Still, it would be males who primarily choose their mating partners. The pattern is male driven, thus cannot be explained by the “extended mating effort hypothesis” (which assumes that the pattern is a product of female choice for mating partners: Smuts & Gubernick, 1992). In accordance with the “extended mating effort hypothesis”, however, we conclude that the relationship between mating and subsequent care does not necessarily indicate paternal investment. Male-female relationships may mediate both, the relationship between infant care and subsequent mating as well as between mating and subsequent care, making it difficult to distinguish between the mating effort and paternal investment hypotheses. These stable relationships may also help males to reduce paternity confusion and increase personal mating success with the female friends at the same time.

Enduring male-female relationship may also result in stability in male infant care, i.e. males taking care of consecutive offspring of the same females. Specifically, in Barbary macaques this stability may be beneficial in relation to the high frequency of male-infant-male interactions, during which two males manipulate one infant together and which help males to manage relationships with other males (Kalbitz et al., 2017; Kuběnová et al., 2019; Paul et al., 1996, Taub et al. 1980). The stable pattern may reduce competition over infants used in these interactions (Kalbitz et al., 2017; Paul et al., 1996), and make those interactions more effective tools for male relationship management (Kubenova et al., 2017; Ogawa, 1995; Paul et al., 1996).

500 Conclusion

501 Sexual selection theory predicts that males and females employ different strategies
502 to increase their fitness, and that some of these strategies may constrain the fitness of the
503 opposite sex (Gavrilets et al., 2001; Parker, 1979). Among others, females and males may
504 adjust the distribution of mating and the extent of infant care. By investigating the link
505 between male infant care and mating in Barbary macaques, we did not confirm the
506 prediction of the mating effort hypothesis from the female perspective. Instead, females
507 seem to increase paternity uncertainty by mating with many males, regardless of care their
508 infant received from particular males. The promiscuous mating behaviour of females
509 makes it difficult for males to track other's mating behaviour rendering such third party
510 knowledge unreliable. This may explain why males turn to direct information on how
511 they distributed their mating across females when making decisions about how to allocate
512 infant care. Males mated more with mothers of infants they had cared for most in the
513 previous season. An untested possibility is that males establish long-term relationships
514 with mothers of their preferred infants who they also pursue for mating opportunities.

515 Thus, the distribution of care and mating in Barbary macaques may be the
516 outcome of contrasting male and female interests in sexual conflict (such as a male's
517 effort for paternal investment constrained by the female's interest to confuse paternity),
518 cognitive constraints (such as the limited ability to track all matings of a female) and
519 additional not immediately reproductive interests (such as the use of infants for male-
520 infant-male interactions). Future research will benefit from more detailed information of
521 female sexual behaviour indicative of female choice, physiological information on
522 conception probability, and genetic paternity data. Future studies may also pay close

attention to intersexual relationships and the role of infants in choosing male care partners.

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772 TABLE 1

Hypothesis	Predictions	Predictors [†]	Response	Supported
Paternal investment hypothesis, female perspective	1) The more a female mated with a specific male (relative to all her matings) the more care her infant will subsequently receive from this male (relative to all male care the infant receives)	1. Distribution of female matings 2. <i>Male rank</i>	Distribution of male care CSI _i [‡] in the birth season following the mating season	No
Paternal investment hypothesis, male perspective	2) The more a male mated with a specific female (relative to all his matings), the more he will subsequently care of her infant (relative to all his infant care)	1. Distribution of male matings 2. <i>Female rank</i>	Distribution of male care CSI _m [§] in the birth season following the mating season	Yes

[†] **Italics: control predictors**

[‡] **Composite social index CSI_i quantifies how much care the infant received from a particular male relative to all other males**

[§] **Composite social index CSI_m quantifies how much a particular male cared for a particular infant relative to all other infants**

Mating effort hypothesis, female perspective	3) The more care an infant received from a specific male (relative to all male care the infant receives), the more often the infant's mother will mate with him (relative to all her matings)	1. Distribution of male infant care in the birth season prior to the mating season 2. <i>Male rank</i>	Distribution of female matings	No
Mating effort hypothesis, male perspective	4) The more a male cared for a specific infant (relative to all his infant care), the more he will subsequently mate with the infant's mother (relative to all his matings)	1. Distribution of male infant care CSIm ^s in the birth season prior to the mating season 2. <i>Female rank</i>	Distribution of male matings	Yes

774 FIGURE LEGENDS

775 *Figure 1. Paternal investment hypothesis based on the female (F) and male (M)*
776 *perspective. Effect of mating on infant care in the subsequent birth season when adding*
777 *to the null model (significant from the male perspective).*

778 *Figure 2. Mating effort hypothesis based on the female (F) and male (M) perspective.*
779 *Effect of infant care in the preceding birth season on the distribution of matings when*
780 *adding to the null model (significant from the male perspective).*

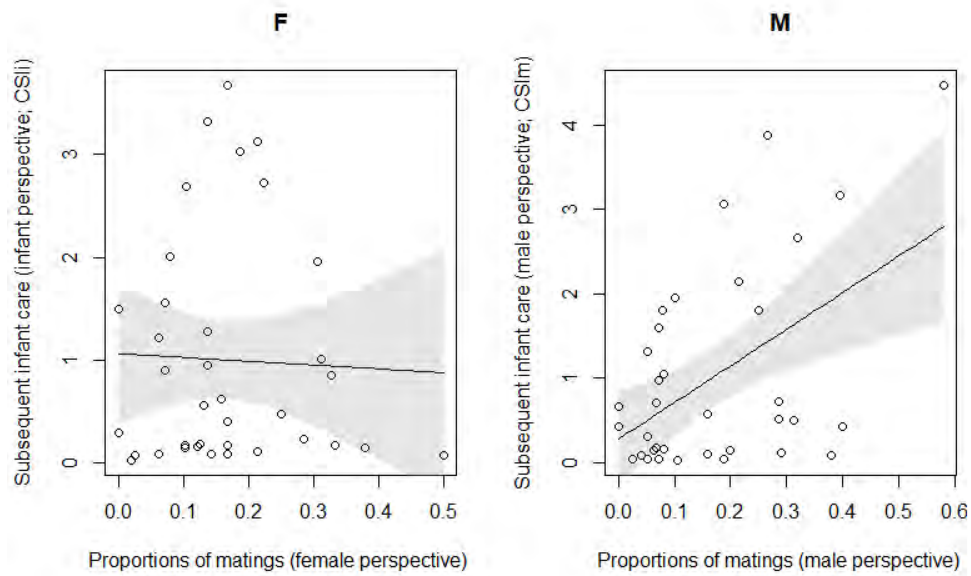


Figure 1. Paternal investment hypothesis based on the female (F) and male (M) perspective. Effect of mating on infant care in the subsequent birth season when adding to the null model (significant from the male perspective).

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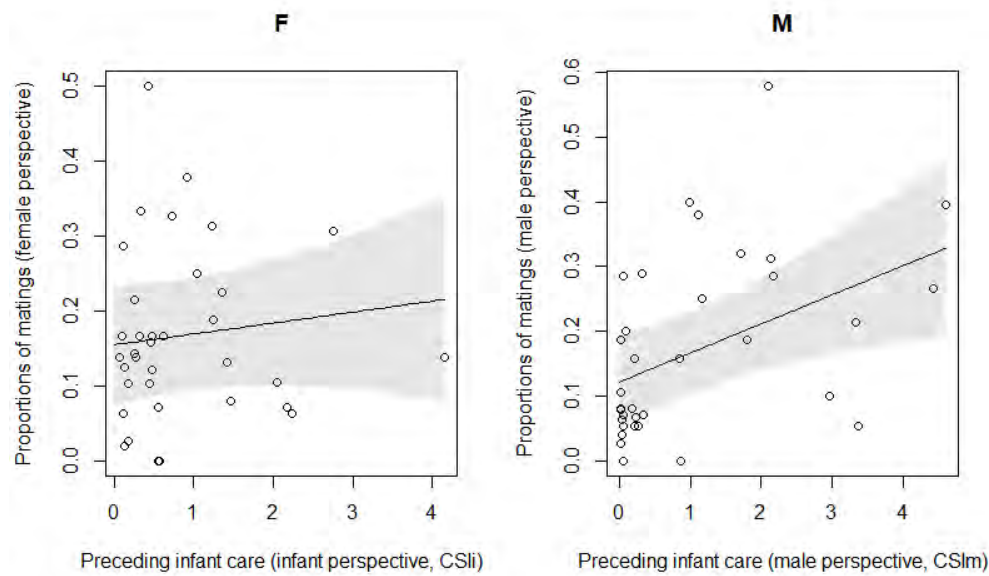


Figure 2. Mating effort hypothesis based on the female (F) and male (M) perspective. Effect of infant care in the preceding birth season on the distribution of matings when adding to the null model (significant from the male perspective).

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